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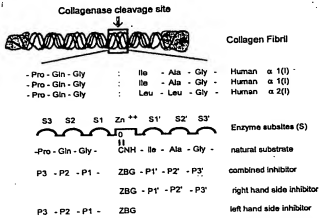


Figure 5. Design of matrix metalloproteinase inhibitors based on the sequence of the collagen substrate cleavage site.

attention<sup>121–123</sup> as right-hand side MMP inhibitors. The early investigations resulted in the identification of left-hand side hydroxamates with only micromolar inhibitory activity such as Z-Pro-Leu-Ala-NHOH (2).<sup>119,120</sup>

A key question that medicinal chemists working in the area have tried to answer is "which ZBG is best"? This issue has been addressed by comparing different ZBGs while keeping the rest of the inhibitor structure constant. Using this approach, Castelhano and co-workers arrived at the following preference in terms of inhibition of MMP-1: hydroxamate (e.g., 3) > formylhydroxylamine > sulfhydryl > phosphinate > aminocarboxylate > carboxylate. 124 Comparison of X-ray crystal structures of 3 and its corresponding carboxylate and sulfodiimine analogues bound to MMP-7 emphasizes the dominant role the ZBG plays in determining the inhibitory potency. The geometries of various ZBGs have been reviewed previously.<sup>76</sup> The hydroxamate acts as a bidentate ligand with each oxygen an optimal distance (1.9-2.3 Å) from the active-site zinc(II) ion, and the position of the hydroxamate nitrogen suggests that it is protonated and forms a hydrogen bond with a carbonyl oxygen of the enzyme backbone. As will become apparent from the discussion below, optimization of the inhibitor structure for many of the above ZBGs can lead to nanomolar inhibition of selected MMPs.

We consider that there are four broad classes of MMP inhibitors as follows: (a) Succinyl hydroxamates, (b) Sulfonamide hydroxamates and related structures, (c) Non-hydroxamates, and (d) Natural products and their derivatives.

While this classification is somewhat arbitrary, it does, we feel, reflect the structural classes of MMP inhibitors that have been investigated. Each of these classes are discussed in turn below with particular emphasis on more recent results.

#### A. Succinyl Hydroxamates

Early studies conducted by Johnson and co-workers demonstrated that succinyl hydroxamic acid derivatives (e.g., 4) are more potent inhibitors of MMP-1 than either the corresponding malonyl (e.g., 5) or

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glutaryl derivatives (e.g., 6).<sup>114</sup> The insertion of a single methylene spacer between the ZBG and the carbon bearing the P1' substituent also showed an improvement in activity for other ZBGs (thiol, formylhydroxylamine, and phosphonate) investigated.<sup>114</sup>

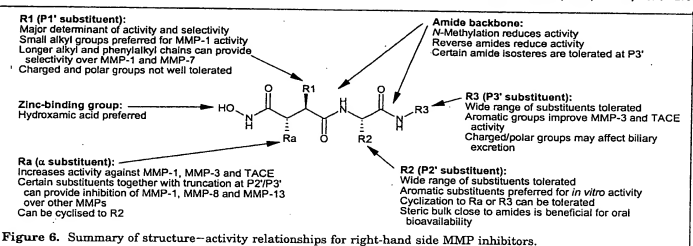
Interestingly, while X-ray analyses of enzyme/ succinyl hydroxamate inhibitor complexes have revealed substrate-like binding modes, X-ray analysis of a malonyl hydroxamate bound to MMP-8 reveals a nonsubstrate-like binding mode. <sup>83</sup> It was discovered that the peptidic tail of the weak inhibitor (2S)-HONH-Mal(i-Bu)-Ala-Gly-NH<sub>2</sub> (7) bound into the deep S1' pocket of MMP-8. <sup>83</sup> This insight has led to the discovery of more potent nonpeptidic malonyl hydroxamates (e.g., 8), <sup>125-127</sup> but the inhibitory activity of these compounds is not as great as that generally obtained with succinyl hydroxamates. The succinyl hydroxamates may be subdivided into peptidic derivatives that possess a P2' amino acid residue and non-peptidyl compounds in which this group is not an amino acid.

## 1. Succinyl Hydroxamates with a P2' Amino Acid Residue

Independently of Johnson and co-workers, <sup>114</sup> the group of Dickens also disclosed that succinyl hydroxamates, such as SC-44463 (9), possess potent MMP inhibitory activity. <sup>128</sup> The structure—activity relationships (SAR) for succinyl hydroxamates possessing a P2′ amino acid residue have been extensively explored. An overview of the SAR is given in Figure 6 and discussed in more detail below.

a. P1 Modifications. We and others have found that generally the introduction of a P1 substituent (substituent a to hydroxamic acid) confers the property of broad-spectrum activity against a variety of MMP enzymes. 129 A beneficial effect is conferred by both lipophilic substituents and those capable of undergoing hydrogen bonding. A P1 alkylcyclic imido substituent increases the potency of not only succinyl hydroxamate MMP inhibitors but also MMP inhibitors with amino carboxylate, phosphinate, and phosphonic acid ZBGs. The role of the cyclic imido group is not entirely clear. In MMP-1, analysis of an X-ray crystal structure has shown that the S1 asparagine (Asn-180; MMP-1 numbering) residue hydrogen bonds to the carbamate carbonyl group of an amino carboxylate inhibitor that possesses a P1 benzyloxycarbonylaminoalkyl substituent.77 A similar interaction is observed between the carbonyl of a P1 phthalimido methyl substituent and Asn-180 in the X-ray structure of Ro 32-0554 (10) complexed to MMP-1 catalytic domain. 130 However, a beneficial effect of a P1' substituent is often observed for the inhibition of MMP-3 even though the corresponding residue to the asparagine is valine which cannot partake in such a hydrogen bonding interaction.

In our own research program, investigation of P1 substituents led to the discovery of batimastat (BB-94) (1),  $^{131,132}$  BB-1101 (11),  $^{133}$  and later marimastat (BB-2516) (12).  $^{129,134}$  Batimastat possesses a thienylthiomethylene  $\alpha$ -substituent and BB-1101 features a smaller allyl  $\alpha$ -substituent, while the  $\alpha$ -substituent for marimastat is a hydroxyl group. All three compounds are broad-spectrum inhibitors which have



displayed efficacy in animal models of human disease' (vide infra). Unlike marimastat, batimastat and BB-1101 are not orally available. We have attributed the oral availability of marimastat in part to the increase in aqueous solubility achieved by the introduction of the \alpha-hydroxyl group. An X-ray crystal structure of BB-1909 (13), an analogue of marimastat, complexed to the catalytic domain of human neutrophil collagenase reveals that the hydroxyl is directed away from the protein surface and is hydrogen-bonded to a solvent molecule. 135 We later found that the presence of certain α-substituents such as allyl (as in BB-1101) and thienylsulfonylmethylene (as in BB-3103 (14)) had a beneficial effect on the inhibition of TACE. This dual activity may be of benefit in diseases which involve both inflammation and matrix remodeling and has been implicated in the pharmacological activity of succinyl hydroxamate compounds, such as BB-1101 (11) in animal models of arthritis 136 and multiple sclerosis.137

Recently, analogues of marimastat have been reported in which the α-position is disubstituted, e.g., 15.138 These compounds feature an α-hydroxy group and an α-methyl group and there is a strong stereochemical preference at the a-position, which is opposite to that of marimastat with respect to the orientation of the hydroxyl. 138 Furthermore, phenylpropyl is reported to be the optimal substituent at P1' and, rather surprisingly, provides potent inhibition of both the short pocket enzyme MMP-1 and the deep pocket enzymes MMP-3 and MMP-9. An X-ray crystal structure of 15 complexed to the catalytic domain of MMP-3 reveals a hydrogen bond between the α-hydroxy group and the backbone of Ala-165 (MMP-3 numbering), as predicted by modeling, and also a van der Waals (VDW) interaction between the P1' aryl group and His-201. The preparation of the succinate portion of these compounds has been published by the Evans group and involves the catalytic asymmetric aldol reaction between methyl pyruvate and the appropriate enolsilane. 139 Gemdisubstitution at the a-position has also been reported for analogues of BB-1101 (11), 140 The combination of  $\alpha$ -methyl and  $\alpha$ -allyl with S stereochemistry (e.g., 16) is well tolerated, whereas activity is reduced when the stereochemistry at the  $\alpha$ -position is R and by gem-diallyl substition. 140 The pharmacokinetics

were investigated for compound 16, but it was found that the quarternary  $\alpha$ -carbon did not confer any benefit compared to BB-1101. We have replaced the  $\alpha$ -hydroxy group of marimastat, respectively, with an  $\alpha$ -alkoxy, e.g., 17, <sup>141</sup> and an  $\alpha$ -cycloalkyl group, e.g., 18. <sup>142</sup> Compound 18 is orally available in the rat and the marmoset and inhibits TNF- $\alpha$  production following oral administration in a rat lipopolysaccharide (LPS) challenge model. <sup>142</sup> The 2,3-disubstituted succinate of compound 18 was prepared by a stereoselective Ireland—Claisen rearrangement. <sup>142,143</sup> The synthesis by solid-phase methods of marimastat analogues in which the  $\alpha$ -hydroxy group is replaced by a substituted  $\alpha$ -amino group has been recently reported. <sup>144</sup>

From analysis of the X-ray crystal structure of batimastat complexed to the active site of recombinant MMP-8 catalytic domain, it is apparent that the  $\alpha\text{-thienylthiomethylene}$  substituent points away from the enzyme as does the P2' phenylalanine side chain.84 Similarly, it has been observed in the X-ray structure of BB-16 (19) complexed to MMP-3 catalytic domain that the P1 and P2' substituents are directed away from the active site into solvent.145 These observations suggest the possibility of joining the P1 and P2' side chains together to form a cyclic inhibitor. Similar cyclication strategies have been followed by Xue and co-workers145 and by Steinman and coworkers. 146 Both groups identified the same compound, SE205 (20), as possessing similar potency to uncyclized analogues. Interestingly, this cyclization strategy resulted in a substantial increase in aqueous solubility (SE205 13 mg/mL vs BB-16 0.3 mg.ml). 145 Increasing the ring size by insertion of one or two methylenes in the alkyl chain from the α-position was well tolerated. 146 Similar inhibitory activity was obtained for the 13-member amide-linked derivative SC903 (21).145 Alternative P1 to P2' macrocyclization strategies have been reported that involve amine formation. 147,148 Depending on the nature of the macrocyclic amine, a degree of selectivity can be obtained for MMP-9 and MMP-8 over MMP-1 and MMP-3147 or activity enhanced against TACE. 148 The introduction of conformational restraint by the construction of a three-membered ring between the a and P1' positions (e.g., 22) has been reported by Martin and co-workers to result in a reduction in the

inhibition of MMP-9. <sup>149</sup> Ghose and co-workers investigated a variety of approaches for the introduction of conformational restraint into the succinyl group in order to determine the pharmacophoric geometry for MMP-1 inhibition. <sup>150</sup> A cyclopropyl derivative (23) that possesses improved in vitro potency in comparison to 22 was identified in this study. <sup>150</sup> The introduction of a six-membered ring between the  $\alpha$  and P1' positions (e.g., 24) resulted in ineffective compounds. <sup>151</sup>

b. P1' Modifications. As discussed earlier, the S1' pocket is considered to be the selectivity pocket for the MMP inhibitors. This is confirmed by SAR data which shows that certain MMPs tolerate large hydrophobic P1' side chains: a P1' 3-phenylpropyl group provides selective inhibition of MMP-2 over MMP-1 and MMP-3 for succinyl hydroxamates (e.g., 25)152 and carboxylates and for phosphonate ZBG MMP inhibitors. 153 This seminal discovery by Porter, Morphy, and co-workers was made before structural data on the MMPs revealed that the S1' subsite is a deep pocket for the majority of the enzymes (e.g., MMP-2, MMP-3, MMP-8, etc.) but is occluded for a few of the MMPs (e.g., MMP-1 and MMP-7) (vide supra). It is intriguing that compound 25, one of the first deep pocket selective MMP inhibitors, should show greater potency for the inhibition of MMP-2 over MMP-3. This tendency for lower IC<sub>50</sub> values against MMP-2 (and the other gelatinase MMP-9) than MMP-3 is exhibited by the majority of MMP inhibitors with extended P1' groups. The discovery that the incorporation of extended P1' groups can provide potent deep pocket selective MMP inhibitors has been embraced and enhanced by medicinal chemists working in this field. An extended alkyl group at P1' provides deep pocket selectivity. Broadhurst and co-workers showed that a C<sub>9</sub> alkyl chain at P1', as in compound 26, gives reduced in vitro inhibition of MMP-1 while maintaining potent inhibitory activity against MMP-2, MMP-3, and MMP-9.154 For a series of matlystatin derivatives, a C9 alkyl chain at P1', as in R-94138 (27), provides at least 10fold greater potency than C<sub>8</sub> or C<sub>10</sub> for the inhibition of MMP-9.155 In analogous succinyl hydroxamates featuring a n-nonyl P1' substituent, the C9 chain length provides at least a 500-fold selectivity for MMP-2 inhibition over inhibition of MMP-1<sup>107,154</sup> yet extending the P1' substituent to C10, as in compound 28, results in potent inhibition of MMP-1.107 Increasing the length of the P1' side chain further to  $C_{16}$ , as in compound 29, results in a loss of activity against MMP-1.107 A similar switch in the inhibition of MMP-1 has been observed within a series of succinyl hydroxamate MMP inhibitors with extended P1' substituents containing heteroatom-based modifications. 156-158 The benzyl ether 30 is a weak MMP-1 inhibitor, whereas the corresponding phenyl ether 31 is a potent MMP-1 inhibitor. 156-158 These results indicate that selected extended P1' substituents can be accommodated in the S1' pocket of MMP-1. In the case of compound 31 it has been proposed that the S1' blocking residue of MMP-1, Arg-214 (MMP-1 numbering), might be displaced by a  $\pi - \pi$  interaction between the electron-rich phenolic group and the

electron-poor guanidinium group. <sup>158</sup> Thus, the occlusion of the S1' pocket for MMP-1 (and presumably that for MMP-7 and MMP-11) is not absolute since the pocket can undergo conformational changes to accommodate certain extended P1' substituents. <sup>76,91</sup> Interestingly, disubstitution at the α-position has been observed to increase MMP-1 potency for a P1' 3-phenylpropyl compound (15)<sup>138</sup> in comparison to a des-α analogue (e.g., 25). <sup>152</sup> Biphenylalkyl P1' substituents have been incorporated into MMP inhibitors with amino carboxyl and carboxylic acid ZBGs. <sup>76,159,160</sup> This modification has also been successfully applied to succinyl hydroxamate compounds (e.g., 32). Similar deep pocket selectivity is observed for MMP inhibitors that feature the related rigid arylalkynylmethylene P1' substituents as in compound 33. <sup>161</sup>

In a series of α-unsubstituted succinyl hydroxamic acid derivatives, phenyl, benzyl, or 2-naphthylmethyl are P1' substituents of choice for the inhibition of soluble CD23 formation. 162 Selectivity for the inhibition of soluble CD23 formation over inhibition of MMP-1 has been achieved by the combination of P1' benzyl with an oxime group at P1 as in compound 34.163 P1' phenyl substitution has also been reported for succinyl hydroxamic acid MMP inhibitors by Robinson and co-workers. 164 They also investigated P1' C-a gem-disubstitution and found that this modification led to a loss of potency relative to the corresponding P1' isobutyl compounds with the least detrimental effect being observed for a P1' gemcyclohexyl compound 35.164 However, a P1' quaternary carbon is tolerated when one of the substituents is hydroxyl as in compound 36165 or when one of the substituents is cyclized onto the nitrogen of the P1'-P2' amide as in compound 37.166

Replacement of the P1'-P2' amide bond of succinyl hydroxamic acid MMP inhibitors by a sulfonamide bond, as in compound 38, results in a substantial loss of MMP inhibitory activity. 167 This has been explained in terms of the hydrogen bond from the N-H of the conserved leucine (Leu-160 for MMP-8) to the sulfonyl oxygen being less energetically accessible due to the pyramidal nature of the sulfonamide. 167 The P1'-P2' amide has also been replaced by a urea functionality, but these analogues were found to be unstable and prone to acid-catalyzed hydantoin formation. 167

c. P2' Modifications. X-ray crystallographic analysis of MMP-inhibitor complexes reveals that the P2' group of peptidyl succinyl hydroxamic acid based MMP inhibitors points out of the enzyme, making few contacts with the S2' cleft. 76 Indeed, analysis of SAR indicates that modification of the P2' group has, in general, a modest effect on in vitro activity. Tryptophan at P2', as in GM6001 (39), 168,169 yields more potent inhibitors than other amino acid side chains. 170 The group at P2' can, however, have an effect on the pharmacokinetic properties of the inhibitors. We have previously suggested that the oral activity of marimastat results from the beneficial combination of a sterically bulky tert-butyl group and an α-hydroxy group which increases aqueous solubility. 129 We argued that the bulky P2' group shields the adjacent amide bonds reducing hydration<sup>171</sup> and, hence, the desolvation energy barrier of the peptide backbone associated with absorption from an aqueous environment to the lipid environment of cell membranes. 172 A P2' tert-butyl group is also a feature of the orally available compounds Ro 31-9790 (40)173 and CT1746 (41).174,175 The phamacokinetics of Ro 31-9790 has been studied in man, and it has been found that the main metabolite is the amide which arises from dehydroxylation of the hydroxamic acid moiety. 176 The deep pocket selective compound CT1746 has been shown to be effective in animal models of cancer following oral administration. 174,175 Babine and Bender suggest that the P2' tert-butyl group is probably inferior with respect to VDW interactions compared to other, more extended side chains but that this is offset by more facile desolvation of the adjacent peptide linkages and a conformational effect that preorganizes the compound for binding.76 Ikeda and co-workers describe compounds that feature a P2' phenyl substituent, e.g., KB-R7785 (42).177 KB-R7785 is orally active as determined both by an ex vivo MMP-1 inhibition assay in mice and demonstration of efficacy in a rat adjuvant arthritis assay. The beneficial effect of the P2' phenyl group on absorption is attributed to it being an amide shielding moiety.177 The same researchers have also investigated the SAR for the inhibition of an MMP-14 mutant lacking the transmembrane domain. 178 It was found that the phenylglycine derivative 42 was a weaker inhibitor of MMP-14 than the corresponding cyclohexylglycine derivative 43, BB-94 (1), BB-2516 (12), or Ro 31-9790 (40). A homology model of MMP-14 suggests that the S1 and S2' subsites are narrower than those of other MMPs. 178 This results in the phenylglycine compound 42 binding in a conformer which is not at an energy minimum whereas the cyclohexylglycine compound 43 and the tert-butylglycine compounds 12 and 14 can bind to MMP-14 in a low-energy comformation.178

The P2' and P3' substituents may be cyclized to form a lactam, and it is found that there is a correlation between the inhibitory potency and ring size for compounds with both hydroxamic acid and phosphonic acid ZBGs. 114,179 This has been attributed to trans amide geometry for the P2'-P3' being the required geometry for effective hydrogen bonding interactions between enzyme and inhibitor. The inhibitory potency was found to increase as the lactam ring size was increased from 7 to 9 and then to 13 atoms. 114 Trans geometry is observed in the complex between MMP-8 catalytic domain and BB-1909 (13), which features a 13-membered lactam ring between P2' and P3'.135 Indolactam cyclization between P2' and P3' (e.g., 3) also confers trans amide geometry and increases activity by at least 10-fold relative to the acyclic analogues. 124

d. P3' Modifications. The S3' region of the MMP enzymes is a relatively open area and a wide range of groups may be introduced at P3'. Heteroaryl and aryl groups appear to enhance MMP-3 and TACE inhibitory activity. We found that the introduction of a benzhydryl group at P3', e.g., 44, leads to compounds that are selective for MMP-7 and MMP-3 relative to MMP-1 and MMP-2. 180,181 Disubstitution

of the P2'-P3' amide tends to reduce inhibitory activity, <sup>114,179</sup> except for P2'-P3' caprolactam derivatives. <sup>182</sup> It has been found that the weak inhibitory activity of P2'-P3' caprolactam derivatives unsubstituted at P3'<sup>114</sup> may be increased substantially by the introduction of a P3' methyl acetate substituent as in compound 45. <sup>182</sup> Interestingly, this compound is a selective inhibitor of MMP-1 over MMP-3 despite possessing an extended C<sub>8</sub> alkyl substituent at P1'. Removal of the P3' ester carbonyl to provide the corresponding methyl ethyl ether results in a reversal of selectivity for inhibition of MMP-3 over that of MMP-1. <sup>182</sup> A P2' dihydrocarbostyril derivative, OPB-3206 (46), exhibits weak MMP inhibitory activity but is orally available in the rat. <sup>183</sup>

The P2'-P3' amide is not strictly required for inhibitory activity since it may be replaced by a variety of alternative functionalities. For example, the P2' amino acid may by replaced by a  $\beta$ -amino alcohol, as in compound 32, but this generally results in a 10-50-fold loss of activity. From the X-ray crystallographic analysis of 32 complexed to the catalytic domain of MMP-3, it is apparent that the C-terminal hydroxy group accepts a hydrogen bond from the N-H of Tyr-240 (MMP-3 numbering).

The replacement of the C-terminal amide group with a nitrogen heterocycle has been a successful modification. An X-ray crystal structure of a Cterminal imidazole 47 complexed to MMP-7 has been reported. 76,184 The use of imidazole to replace the C-terminal N-methyl amide is said to result in a 5-fold reduction in potency against all MMPs tested for this analogue of GM6001.76 Examination of the crystal structure shows that the imidazole makes hydrogen bonds to the carbonyl of Asn-179 and to the N-H of Tyr-240 (MMP-7 numbering) without any apparent perturbation of the inhibitor backbone in comparison to a related structure in which the C-terminal N-methyl amide is retained. 76,184 The introduction of a phenyl substituent at the 5-position of the imidazole ring provided selective inhibition of MMP-7 over MMP-1 and MMP-3, whereas a P3' benzimidazole group provided broad-spectrum inhibition of the three MMPs tested. 184

The replacement of the P2'-P3' amide by an aryl ketone or heteroaryl ketone group, as in the P3' indole ketone 48 and the P1-P2' cyclized P3' phenyl ketone 49, is tolerated. 185 Compound 48 is an analogue of BB-1101 (11) which possesses negligible oral availability. In contrast, the indole ketone 48 is 12% bioavailable in the monkey following dosing at 10 mg/ kg p.o. and has a half-life of 20 h.185 Two earlier studies of P2'-P3' amide derivatives showed that the nature of the P3' substituent can have an effect on the oral availability of succinyl hydroxamates. 186,187 In one study, the effect of P3' substituents on biliary excretion in the rat was examined for a series of GM6001 analogues. 186 It was found that the presence of a tertiary amine at P3' reduced biliary excretion and increased plasma half-life. 186 The tertiary amine 50 was found to be 8.5% orally bioavailable in the rat. 186 In the other study, it was found that oral availability as measured in a mouse pleural cavity assay was significantly enhanced by the introduction of an alkyl morpholino P3' substituent.<sup>187</sup> The beneficial effect of the morpholino group was attributed to its basicity.<sup>187</sup> Epimerisation-free amide coupling conditions have been developed by Fray and Ellis to facilitate the introduction of a wide range of P3' substituents into a N-succinyl-tert-leucine intermediate for the preparation of succinyl hydroxamates.<sup>188</sup>

### **B. Non-Peptidic Succinyl Hydroxamates**

Truncation of the P2'-P3' group of pseudo-peptide succinyl hydroxamic acid derivatives leads to MMP inhibitors which tend to be selective for the collagenases. Broadhurst and co-workers discovered that potent inhibition of the collagenases could be achieved when a cyclic imide group is introduced at P1, as in the phthalimido derivative 51.189 Subsequent optimization of this series led to the discovery of Ro 32-3555 (52), which was selected for development for the treatment of arthritis. 190,191 Presumably, for compound 51 and Ro 32-3555 (52), a favorable balance between active-site interactions and solvation is maintained despite the removal of three hydrogen bonding groups in comparison to succinyl hydroxamates with P2' groups. The presence of the cyclic imide group at P1, a feature of earlier MMP inhibitors identified by the same workers and by other groups (vide supra), appears to also be important for activity and may compensate for the loss of the hydrogen bonds as observed in the X-ray structure of Ro 32-0554 (10) complexed to the active site of MMP-1.130 The cyclopentylmethyl P1' group of Ro 32-3555 was chosen on the basis of X-ray crystal structure data for MMP-1. 190 The introduction of cyclopentylmethyl provides a modest increase in potency over an analogue of Ro 32-3555 with an isobutyl group at P1' suggesting improved complementarity with the S1' pocket. 190 Ro 32-3555 exhibits an oral bioavailability of 26% in the rat and inhibits articular cartilage degradation in a rat monoarthritis model. 192 Ro 32-3555 (Trocade) has been referred to as a cartilage protective agent (CPA). 193 An improved synthesis of the chiral 2,3-disubstituted succinate of Ro 32-3555 has been reported 194 based on the earlier succinate alkylation protocol of Crimmin and coworkers. 195 We found that incorporation of a sulfonamide moiety at P1 in combination with P2'-P3' truncation, as in compound 53, can provide selective inhibition of MMP-1 over other the MMPs that we evaluated. 196 Analogues of 53 possess different selectivity profiles depending on the nature of the sulfonyl substituent.

Alpegiani and co-workers found that an α amino group in conjunction with a P2' piperazinyl moiety, as in compound 54, provided compounds with good oral availability. 197 Compound 54 exhibits 58% oral bioavailability in the rat and 34% in the cynomolgus monkey. 197 Broadhurst and co-workers have found that the P2'-P3' amino acid residue may be replaced by a hydrazide moiety as in Ro 32-7315 (55). 198,199 This potent TACE selective inhibitor has been selected for clinical development. 199 Cyclic hydrazide compounds such as the piperazic acid derivative matlystatin B (56) and its analogues have been previously identified as natural product MMP inhibi-

tors. 155,200-202 Matlystatin analogues have been prepared in which the ZBG has been altered, 203 the P3' group modified, 155,204 and the P1' substituent changed. 155 As described above, the P1' nonyl derivative 27 is a potent inhibitor of the gelatinases, in contrast to the modest activity exhibited by the parent molecule matlystatin B (56). 155

The P2' amino acid residue of succinyl hydroxamic acid MMP inhibitors may be replaced with a benzhydryl group, as in compound 57.111 A combinatorial chemistry approach was employed in this study that first involved the exploration of P2' modifications for a series of N-carboxyalkyl amino acid based inhibitors (vide infra). Modeling of the optimal P2' group that had been identified by this approach led to the identification of benzhydryl as a preferred substituent, which was then introduced into the corresponding succinyl hydroxamic acid derivative 57. An X-ray crystal structure of this compound bound to the catalytic domain of MMP-3 revealed an unexpected conformational shift in the 222-231 loop region (MMP-3 numbering).111 This illustrates that subtle changes in binding can occur with variation of inhibitor structure and that these are very difficult to predict on the basis of modeling alone. Inhibition of MMP-3 was further increased by replacing one of the phenyl groups of compound 57 with 3-pyridyl. 111

# C. Sulfonamide Hydroxamates and Related Structures

N-Sulfonyl amino acid hydroxamates were independently identified as inhibitors of MMPs by two research groups. 205,206 The first such compound to enter development is the orally available broadspectrum inhibitor CGS 27023A (58).205 Key structural features of CGS 27023A are said to be the isopropyl substituent which slows down metabolism of the adjacent hydroxamic acid group and the basic 3-pyridyl substituent which may aid partitioning into the hydrated negatively charged environment of cartilage.205 SAR for the inhibition of macrophage metalloelastase (MMP-12) by CGS 27023A and analogues has been reported.207 This reveals that CGS 27023A is a potent inhibitor of MMP-12, an enzyme that has been implicated in the development of emphysema that results from chronic inhalation of cigarette smoke. 208 The binding mode of CGS 27023A and analogues to MMP-3 has been investigated by NMR spectroscopy. 209-211 The p-methoxyphenyl substituent of CGS 27023A occupies, but does not fill, the S1' specificity pocket, while the pyridylmethyl and isobutyl substituents occupy the S2' and S1' subsites, respectively.211 An X-ray crystallographic analysis of a related compound CGS 25966 (59), a close analogue of CGS 27032A, complexed to the catalytic domain of MMP-1 has been reported by Babine and Bender. 76 The observed binding mode is broadly in agreement with the NMR studies in that the 4-methoxyphenyl group resides in the S1' pocket and the isopropyl group is located in the S1 subsite. The X-ray structure indicates that the isopropyl group is relatively close to the N-benzyl substituent.76 By introducing a six-membered ring to provide beneficial ligand preorganization76 and extending the P1' substituent, the potent MMP inhibitor AG3340 (60) was derived. 212,213 This compound was selected for development based on its superior efficacy in a murine model of cancer growth and metastasis in comparison to a number of analogues and because it showed a favorable pharmacokinetic profile with 18% oral bioavailability in rats.<sup>213</sup> There has been considerable interest from other researchers in analogues of CGS 27023A and AG3340.141 For example, it has been recently reported by Hanessian and co-workers that modification of the substituent a to the hydroxamic acid in CGS 27023A leads to increases in the inhibition of the deep pocket MMPs, e.g., thioether derivative 61.<sup>214</sup> Other analogues of particular interest are sulfone derivatives, <sup>215–218</sup> bis-sulfonamides, <sup>219,220</sup> and phosphinamides.<sup>221</sup> Groneberg, Burns, and co-workers have identified sulfone hydroxamic acids (e.g., 62) which are inhibitors of both MMPs and the enzyme phosphodiesterase type 4 (PDE4).216 Inhibition of PDE4 results in increased intracellular concentration of cyclic AMP and consequently in antiinflammatory activity.222 Analogues of compound 62 have been identified, by both combinatorial methods (vide infra)215 and traditional analogue synthesis,216 that provide selective inhibition of PDE4 over MMP inhibition by the introduction of a 3,4-dimethoxyphenylsulfonyl group. The reverse selectivity for MMP inhibition over that of PDE4 is achieved by the incorporation of a cyclic quaternary center a to the sulfonyl moiety.215 This is a structural feature of the compound RS-113,456 (63) identified by Campbell and co-workers. 217,218 Oral availability and half-life were improved in this series by shifting the cyclic group to be a to the hydroxamic acid as in the development compound RS-130,830 (64).217 Separate X-ray crystallographic analyses of both RS-113,456 and RS-130,830 bound to the catalytic domain of MMP-13 reveal that the two compounds adopt virtually identical conformations.91 An X-ray crystal structure has also been determined for RS-104,966, an analogue of 63 lacking the chloro substituent, bound to the catalytic domain of MMP-1. This shows that induced fit of MMP inhibitors with large P1' substituents can occur by Arg-214 (MMP-1 numbering) adopting a new position, creating a larger open S1' pocket.91 RS-113,-456 (63) dosed orally diminishes flow-mediated arterial enlargement in a rat arteriovenous fistula model,217 and RS-130,830 (64) is being investigated in the clinic as a therapeutic agent for the treatment of osteoarthritis.218 In contrast to the majority of MMP inhibitors, both RS-113,456 and RS-130,830 lack any stereocenters yet retain potent inhibitory activity for the deep pocket MMPs. In an alternative approach based on symmetrical bis-sulfonamides, Pikul and co-workers identified another series of nonchiral MMP inhibitors (e.g., 65).219 The 1,3-piperazinyl derivative PGE-4410186 (65) exhibits broadspectrum inhibitory activity against the enzymes tested.219 An X-ray crystal structure of PGE-4410186 complexed to the catalytic domain of MMP-3 reveals similar interactions previously observed in the crystal structure of CGS 25966 complexed to the catalytic domain of MMP-176 in terms of one of the 4-methoxyphenylsulfonyl groups residing in the S1' pocket.219 The second 4-methoxyphenylsufonyl binds to the S1/ S2 pocket with the two sulfonyl oxygen atoms of this group interacting with the imidazole ring of His-166 (MMP-3 numbering) via a hydrogen-bonded bridging water molecule.219 PGE-4410186 and analogues have been evaluated in an in vitro cartilage permeation model.<sup>220</sup> It was found that permeability across articular cartilage was increased for analogues of PGE-4410186 with increasing hydrophilicity. 220 Pikul and co-workers have also investigated analogues of CGS 27023A in which the sulfonamide moiety is replaced by a phosphinamide group as in compound 66.221 This compound is a potent inhibitor of MMP-3, the collagenases (MMP-1, MMP-8, MMP-13), and the gelatinases (MMP-2, MMP-9) but is less effective at inhibiting MMP-7. An X-ray crystal structure of phosphinamide 66 bound to MMP-3 catalytic domain reveals that the phosphinamide phenyl group is accommodated into the S1' pocket and that the phosphinamide oxygen is within hydrogen bonding distance to the N-H of Leu-164 and Ala-165 (MMP-3 numbering).221 This provides an explanation for the observation that optimum enzyme inhibitory activity is achieved when the configuration at the phosporus chiral center is R.221 However, hydrolysis of the phosphinamide bond which occurs at low pH may limit the potential of these compounds to be developed into orally available drugs.

A new drug discovery technique involving multidimensional NMR spectroscopy, known as "SAR by NMR", was first used to identify potent analogues of the immunosuppressant FK506.<sup>223</sup> Application of this technique has been extended to the field of MMP inhibition, where it has been used to identify a series of MMP-3 inhibitors.<sup>224,225</sup> In this study, two ligands that bind weakly to proximal sites on MMP-3 were identified. Acetohydroxamic acid binds to the activesite zinc(II) ion and 3-cyanomethyl-4'-hydroxybiphenyl binds to the S1' pocket.<sup>224</sup> On the basis of the NMR-derived structural information, the two molecules were linked together to give the potent MMP-3 inhibitor 67.<sup>224</sup>

### D. Non-Hydroxamates

Due to the intense competition in the area of hydroxamic acid MMP inhibitors there has been considerable interest in compounds with alternative zinc binding groups. For the purpose of this review these are subdivided as follows: (1) Carboxylic acid and N-carboxyalkyl ZBGs, (2) Thiol ZBGs, (3) Phosphorus-based ZBGs, (4) Novel zinc binding groups.

## 1. Carboxylic Acids and N-Carboxyalkyl ZBGs

With the exception of a few families of hydroxamic acids derived directly by solid-phase procedures, all the other examples of MMP inhibitors described in the preceding sections will have been prepared by converting a carboxylic precursor into the corresponding hydroxamic acid. Consequently, a vast volume of test data has been built up on the value of carboxylic acid containing structures as potential MMP inhibitors. A typical example is the carboxylic acid precursor of compound 29; this is selective for MMP-2 over MMP-1, -3, and -7 (IC<sub>50</sub> of 30 nM vs

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